Development of Wheat Lines Having a Small Introgressed Segment Carrying Stem Rust Resistance Gene Sr22

Eric L. Olson, Gina Brown-Guedira,* David Marshall, Ellen Stack, Robert L. Bowden, Yue Jin, Matthew Rouse, and Michael O. Pumphrey

ABSTRACT

The wheat stem rust resistance gene Sr22 confers resistance to Puccinia graminis f. sp. tritici Pers. race TTKSK (also known as Ug99) that developed in Africa and is an immediate threat to world wheat production. The resistance gene is present on a chromosomal translocation derived from Triticum boeoticum Boiss., which has a genome that is partially homologous to the A genome of T. aestivum L. Sr22 has been deployed in a limited number of cultivars due to poor agronomic performance of lines carrying the resistance gene. Linkage analysis of simple sequence repeat (SSR) markers on chromosome 7A was performed to identify loci closely linked to Sr22. The most tightly linked proximal and distal SSR marker loci were Xcfa2123 and Xwmc633, respectively. A two-step process was then used to develop resistant lines having smaller chromosome segments derived from the diploid donor. First, individuals in which a single recombination event had occurred between wheat and the Sr22 introgression were identified in the mapping populations. In spite of reduced recombination between T. boeoticum and T. aestivum chromosomes, sufficient recombination events were found among 398 F_{3:4} lines derived from recombinant F₂ progeny to recover multiple resistant individuals with smaller alien introgressions. Resistant lines were identified having less than 6% of the chromosome arm derived from T. boeoticum. These lines may provide a more agronomically desirable source of Sr22 that can be readily deployed in cultivars resistant to Ug99.

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Abbreviations: FL, fraction length; IT, infection type; NIL, near isogenic line; PCR, polymerase chain reaction; SSR, simple sequence repeat.

WILD RELATIVES of modern bread wheat (*Triticum aestivum* L.) are an important source of genes conferring resistance to disease and insect pests. Several genes for resistance to stem rust caused by *Puccinia graminis* f. sp. *tritici* Pers. (*Pgt*) have been introgressed from related species, including the adult plant resistance genes *Sr2* from the *T. turgidum* L. emmer wheat landrace 'Yaroslav' (McFadden, 1930), *Sr24* from tall wheatgrass [*Thinopyrum elongatum* (Host.) D.R. Dewey] (Sears, 1973), *Sr31* from rye (*Secale cereale* L.) (Zeller, 1973), and *Sr36* from timopheevi wheat [*T. timopheevii* (Zhuk.) Zhuk.] (Nyquist, 1962). These resistance genes have been widely deployed

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to provide stem rust resistance in many wheat growing areas of the world. Other broadly effective *Sr* genes transferred from wheat relatives, including *Sr32*, *Sr39*, and *Sr40*, remain undeployed in modern agriculture due to reduced agronomic performance associated with their presence on large alien chromosome introgressions (Singh et al., 2008).

The resistance gene Sr22 was introgressed into common wheat from wild and cultivated A-genome diploid wheats (The, 1973). It provides effective resistance to the variant stem rust races of the TTKS lineage (also referred to as Ug99) (Jin et al., 2007) that developed in Africa and now threaten world wheat production. The Sr22 gene was identified in the T. boeoticum Boiss. [syn. T. monococcum ssp. aegilopoides (Link) Thell.] accession G-21 (Gerechter-Amati et al., 1971) and the T. monococcum L. (syn. T. monococcum L. ssp. monococcum) accession RL5244 (Kerber and Dyck, 1973). Introgressions from the two donors differ in the amount of alien chromatin present. The *T. boeoticum* introgression into the common wheat cultivar Steinwedel includes nearly the entire long arm and a portion of the short arm of chromosome 7A^m. The T. monococcum introgression into Marquis consists of the distal region of 7A^mL (Kerber and Dyck, 1973; Paull et al., 1994). Khan et al. (2005), using simple sequence repeat (SSR) markers, genetically mapped Sr22 between Xcfa2123 (proximal) and Xcfa2019 (distal) with an interval distance of 11.2 cM, an interval previously estimated to be approximately 30 cM (Sourdille et al., 2005). The smaller genetic distance in the Sr22 mapping population was presumably the result of reduced recombination between the introgressed segment carrying the gene and wheat chromosome 7A.

Reduced recombination between chromosomes of the A genome of *T. aestivum* and the A^m genome of einkorn wheat relatives has been well documented (Dubcovsky et al., 1995; Luo et al., 2000). The Ph1 gene of hexaploid wheat restricts homoeologous pairing (Riley and Chapman, 1958), preventing recombination between homoeologous chromosomes. Consequently, introgressed chromosome segments from species that are not completely homologous may be inherited in large linkage blocks carrying substantial amounts of alien chromatin. Detriments to agronomic performance have been reported in association with the Sr22 introgressions, including depression of yields and delayed heading date (Paull et al., 1994). In near isogenic lines (NILs) from different genetic backgrounds carrying Sr22, The et al. (1988) reported a mean yield that was not significantly lower than non-Sr22 controls. However, the results varied among genotypes, with some Sr22 NILs yielding as much as 10% less than non-Sr22 controls. Paull et al. (1994) reported that lines without the T. boeoticum introgression headed approximately 6 d earlier than lines heterozygous and homozygous for the introgression. In addition, reduced transmission of gametes carrying Sr22 has been observed for both the T. boeoticum and T. monococcum introgressions (The and McIntosh, 1975). Poor agronomic performance and reduced transmission of Sr22 have been

attributed to linkage drag resulting from substantial chromatin derived from the A^m genome relatives.

The associated negative effects on agronomic performance have hindered the use of *Sr22* worldwide. However, deployment of this gene is of particular interest in contemporary wheat breeding because it confers resistance to *Pgt* race TTKSK and more virulent derivatives that have emerged from East Africa (Singh et al., 2008). Additionally, *Sr22* confers resistance to all domestic *Pgt* races in the United States. In this study we report genetic linkage analysis of the *Sr22* region in multiple populations and the identification of more tightly linked SSR marker loci for use in marker-assisted selection. In addition, recombinants have been identified that carry reduced amounts of *T. boeoticum* chromatin that may prove useful in the deployment of *Sr22* in breeding populations.

MATERIALS AND METHODS

Plant Materials

For linkage analysis of *Sr22*, two mapping populations were developed. The resistant parent for both crosses was the germplasm Sr22Tb with the pedigree Steinwedel*2//Spelmar*2/*T. boeoticum* G-21. Seed of this Steinwedel selection that has the 7A^m chromosomal translocation from *T. boeoticum* carrying the stem rust resistance gene *Sr22* (The, 1973) was provided by Dr. Yue Jin, USDA-ARS Cereal Disease Lab, St. Paul, MN. Two Steinwedel accessions, PI 41081 and PI 27018, obtained from the USDA National Small Grains Collection were used to compare the haplotypes of SSR loci linked to *Sr22* between the germplasm Sr22Tb and the original cultivar Steinwedel.

A population of 138 F_{2:3} lines, referred to as U5615-F_{2:3}, was developed from the cross of Sr22Tb with the hard winter wheat cultivar 2174 (PI 602595). An additional population of 139 $F_{2:3}$ lines, referred to as U5616- $F_{2:3}$, was developed from a cross between Sr22Tb and the hard winter wheat cultivar Lakin (PI 617032). From these populations, three F_{2:3} families having recombination between distal marker Xcfa2019 and proximal marker Xbarc121 in the Sr22 region were selected for development of additional mapping populations and identification of recombinants having smaller introgression segments. Linkage analysis was performed using SSR loci segregating in F_{3:4} populations developed from the F₂ plants U5615-72, U5615-98, and U5616-20. From the recombinant plant U5615-72 that was homozygous for alleles from parent line 2174 at the distal Xcfa2019 locus and heterozygous for Sr22, 104 F_{3:4} lines were developed (population U5615-72-F_{3.4}). From the U5615-98 recombinant that is homozygous for alleles from parent line 2174 at the proximal Xbarc121 locus and heterozygous for Sr22, 140 F_{3.4} lines were developed (population U5615-98-F_{3:4}). A population of 152 F_{3:4} lines was derived from the recombinant F₂ plant U5616-20 that was homozygous for Lakin alleles at the Xbarc121 locus and heterozygous for Sr22 (population U5616-20-F_{3.4}).

The chromosomal assignment of the SSR loci to the long arm of chromosome 7A was done using Chinese Spring nullisomic-tetrasomic and ditelosomic stocks (Sears, 1954; Sears and Sears, 1978) for this chromosome. Chromosome physical mapping of SSR loci was done using Chinese Spring chromosomal

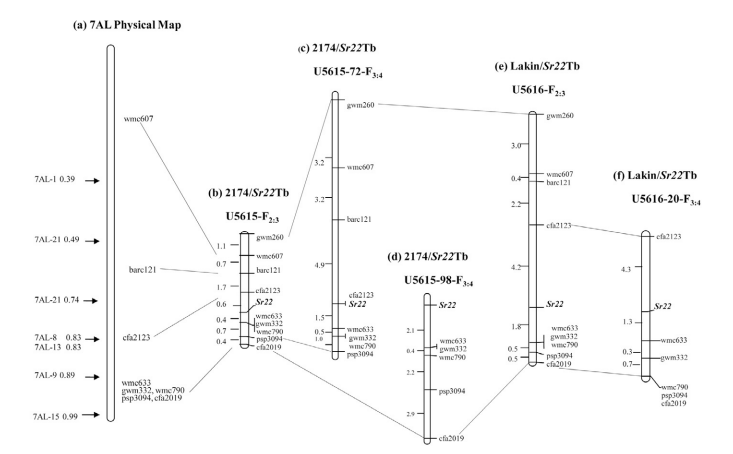


Figure 1. (a) Physical map of SSR loci linked to Sr22 on the long arm of chromosome 7A (Xgwm260 not shown because it is located on 7AS). Names of deletion lines and the deletion breakpoints shown at left. (b–d) Genetic linkage of F $_{2:3}$ population U5615 (2174/Sr22Tb) and F $_{3:4}$ populations U5615-72, U5615-98 derived from recombinant F $_2$ individuals segregating for stem rust resistance from Sr22. (e–f) F $_{2:3}$ population U5616 (Lakin/Sr22Tb) and F $_{3:4}$ population U5616-20 derived from a recombinant F $_2$ individual segregating for stem rust resistance from Sr22.

deletion stocks of 7AL. The lines used in deletion mapping include 7AL-15 (0.99), 7AL-9 (0.89), 7AL-13 (0.83), 7AL-8 (0.83), 7AL-21 (0.74), 7AL-10 (0.49), and 7AL-1 (0.39) (Endo and Gill, 1996) (Fig. 1a). All aneuploid and deletion stocks were obtained from the Wheat Genetics and Genomics Resource Center at Kansas State University, Manhattan, KS.

Stem Rust Evaluations

Puccinia graminins f. sp. tritici, race RKQQC (virulence/aviulence formula: Sr5, 6, 7b, 8a, 9a, 9b, 9d, 9 g, 21, 36, McN/ Sr9e, 10, 11, 13+17, 24, 30, 31, 38, Tmp [based on revised nomenclature from Jin et al., 2008]) is avirulent to Sr22 and was used for phenotypic analysis of mapping populations. At least 10 seedlings of all F_{2:3} and F_{3:4} lines and the resistant and susceptible parents were grown in 10 by 10 cm square pots in Metro-Mix 200 vermiculite-peat-perlite medium (Hummert, Inc., Earth City, MO) in a greenhouse. Urediniospores were removed from liquid nitrogen storage and heat-shocked in a 42°C water bath for 5 min. Spores were suspended in Soltrol 170 isoparaffin oil (Chevron Phillips Chemical Company LP, The Woodlands, TX) and sprayed onto two- to three-leaf stage seedlings. Inoculated plants were incubated in a dew chamber at 24 ± 1 °C, 100% relative humidity for 16 h and then grown in a greenhouse at 21 ± 4°C with 16 h light/8 h dark cycle. Infection types (ITs) described by Stakman et al. (1962) were assessed 14 d after inoculation. Rust evaluations were replicated so that a total of at least 20 seedlings from each line were evaluated.

Stem rust reaction of parent lines and selected recombinants having reduced *T. boeoticum* chromatin were determined using eight races of *P. graminis* f. sp. *tritici*, including races of the TTKS lineage at USDA-ARS Cereal Disease Laboratory (St. Paul, MN). Inoculation and disease assessment procedures were reported previously (Jin et al., 2007). Virulence/avirulence formulas of the tested races are given in Table 1.

Molecular Marker Analyses

Genomic DNA was extracted from F_2 plants and from a bulk of 10 plants of each $F_{3:4}$ line. DNA isolations were done according to a protocol modified from Pallotta et al. (2003) or using a QIAGEN DNeasy 96 Plant kit (QIAGEN, Valencia, CA) according to the manufacturer's instructions.

Parent lines 2174, Lakin, Sr22Tb, and Steinwedel were screened for polymorphism with 42 SSR markers previously mapped to the long arm of chromosome 7A including: barc29, barc49, barc108, barc121, barc174, barc192, barc195, barc221, cfa2019, cfa2040, cfa2123, cfa2257, cfd20, cfd68, cfd193, gwm4, gwm10, gwm63, gwm260, gwm276, gwm282, gwm332, gwm344, gwm473, gwm554, gwm573, psp3094, wmc17, wmc96, wmc116, wmc139, wmc107, wmc139 wmc273, wmc346, wmc426, wmc488, wmc525, wmc607, wmc633, wmc790, and

Table 1. Virulence and avirulence formulas for the Puccinia graminis f. sp. tritici races used in the phenotypic analysis of Sr22.

Pgt race [†]	Avirulence	Virulence
MCCFC (59KS19)	6 8a 9a 9b 9d 9e 11 21 24 30 31 36 38	5 7b 9g 10 17 Tmp McN
QFCSC (06ND76C)	6 7b 9b 9e 11 24 30 31 36 38 Tmp	5 8a 9a 9d 9g 10 17 21 McN
QTHJC (75ND717C)	7b 9a 9e 24 30 31 36 38 Tmp	5 6 8a 9b 9d 9g 10 11 17 21 McN
RKQQC (99KS76A)	9e 10 11 17 24 30 31 38 Tmp	5 6 7b 8a 9a 9b 9d 9g 21 36 McN
TPMKC (74MN1409)	6 9a 9b 24 30 31 38	5 7b 8a 9d 9e 9g 10 11 17 21 36 Tmp McN
TTTTF (01MN84A-1-2)	24 31	5 6 7b 8a 9a 9b 9d 9e 9g 10 11 17 21 30 36 38 McN Tmp
TTKSK (04KEN156/04)	24 36 Tmp	5 6 7b 8a 9a 9b 9d 9e 9g 10 11 17 21 30 31 38 McN
TTKST (06KEN19-V-3)	36 Tmp	5 6 7b 8a 9a 9b 9d 9e 9g 10 11 17 21 24 30 31 38 McN

[†]Pgt race code, after Roelfs and Martens (1988) using revised nomenclature from Jin et al. (2008). Race isolates given in parentheses.

wmc809. Markers amplifying polymorphic fragments were run on aneuploid and deletion stocks to validate their chromosomal location and then evaluated on mapping populations. Thirteen markers amplifying polymorphic fragments between Sr22Tb and both hard winter wheat parents and assigned to chromosome 7A were used for linkage analysis of segregating populations.

The polymerase chain reaction (PCR) master mix for SSR primers consisted of 2 µL of 20 ng µL⁻¹ genomic DNA template, 0.40 µL of a 10 µM mixture of forward and reverse primers, 0.18 μ L (0.9 U) of Taq polymerase, 1.20 μ L of 10× buffer (10 mM Tris-HCl, 50 mM KCl, 1.5 mM MgCl₂, pH 8.3), 0.96 μ L of a 100 μ M mixture of dNTPs, and 7.26 μ L of water, bringing the total reaction volume to 12 μL. A touchdown profile was used that consisted of an initial denaturation at 95°C followed by 15 cycles of 95°C (45 s), 65°C (45 s) decreasing by 1°C each cycle, and 72°C (60 s), followed by 25 cycles of 50°C annealing temperature. The cycling conditions for markers requiring specific annealing temperatures included an initial denaturation of 95°C followed by 35 cycles of 95°C (45 s), 49°C or 60°C (45 s), and 72°C (60 s), followed by a final extension at 72°C (4 min). The forward primers for all SSR markers were 5'-modified to include the fluorescent dye 6-FAM. Amplifications were performed using an Eppendorf Mastercycler (Eppendorf AG, Hamburg, Germany). Sizing of PCR products was performed by capillary electrophoresis using a 3130xl Genetic Analyzer (Applied Biosystems, Foster City, CA). Analysis of PCR fragments was performed using GeneMarker 1.60 software (SoftGenetics, State College, PA). Genetic linkage analysis was performed using MAPMAKER v 3.0 (Lander et al., 1987). Marker order was established using multipoint analysis and the Haldane centimorgan function with a minimum LOD of 3.0. Segregation of marker loci and resistance reaction was evaluated using a χ^2 goodness-of-fit test. Differences in recombination frequency between populations were determined according to Allard (1956).

RESULTS AND DISCUSSION

Phenotypic Evaluation

Germplasm Sr22Tb exhibited ITs of 2- or 2 when inoculated with diverse races of *Pgt*, including races TTKSK and TTKST (Table 2). Susceptible reactions (IT = 4) were observed on parent line 2174 when inoculated with six of the eight races of *Pgt*. Resistant reactions were observed when cultivar 2174 was inoculated with isolates MCCFC and RKQQC, indicating the presence of a race-specific

stem rust resistance gene in this parent. The infection type was variable among different inoculation experiments when 2174 was challenged by RKQQC, suggesting the presence of an environmentally sensitive resistance gene(s) in this line (data not shown). The parent line Lakin was consistently susceptible to all isolates.

In each segregating population, individuals homozygous or heterozygous for *Sr22* exhibited a low IT of 1– to 2 on challenge with *Pgt* race RKQQC. Uredinia were typically small and surrounded by chlorosis rather than necrosis. Individuals in the populations with an IT of 2+ or greater were considered susceptible (*sr22sr22*).

Analysis of segregation ratios indicated that resistance to stem rust was due to a single dominant gene in both populations (Table 3). Although reduced transmission of *Sr22* had been previously reported by Paull et al. (1994), there was no significant deviation from the 1:2:1 segregation ratio expected for a single dominant gene. Encouragingly, the variable seedling resistance to RKQQC in 2174 did not interfere with our ability to score *Sr22* in F₃ families.

Genetic and Physical Mapping of the Sr22Tb Introgression

Of the 42 SSR loci evaluated, 13 were polymorphic between Sr22Tb and the hard winter wheat parents 2174 and Lakin. Linkage analysis of these makers on the U5615-F_{2:3} and the U5616-F_{2:3} populations placed the markers onto two linkage groups. Nine of the SSR markers were placed onto a linkage group with Sr22. These loci included Xbarc121, Xcfa2019, Xcfa2123, Xgwm260, Xgwm332, Xpsp3094, Xwmc607, Xwmc633, and Xwmc790 (Fig. 1). Four other loci, Xcfa2257, Xgwm344, Xwmc346, and Xwmc525, formed a second unlinked group greater than 60 cM distal to Sr22 (LOD > 3; map not shown). These four markers have previously been located terminally on 7AL (Miranda et al., 2007; Perugini et al., 2007).

The orders of markers in the Sr22 linkage group agree in both populations (Fig. 1). The nine linked markers covered a region of 5.6 cM in the U5615-F_{2:3} population, compared to a distance of 12.6 cM in the U5616-F_{2:3} population. Thus, there was a twofold reduction in recombination in the 2174/Sr22Tb population compared with the population derived from the cross Lakin/Sr22Tb (P < 0.01). In

Table 2. Infection types of parent lines 2174, Lakin, Sr22Tb, and selected recombinants when inoculated with eight different races of *Puccinia graminis* f. sp. *tritici*.

Line	TTKSK (04KEN156/04) [†]	TTKST (06KEN19-V-3)	TPMKC (74MN1409)	TTTTF (01MN84A-1-2)	RKQQC (99KS76A)	QFCSC (06ND76C)	QTHJC (75ND717C)	MCCFC (59KS19)
Lakin	4	3	4	4	4	4	-	4
Sr22TB	2-	2-	2	2	2-	;2-§	2	2-
2174	4	4	4	4	3/13-	4	4	;2-
U5615-98-120‡	2	2-/4¶	2/4	2	2-/3	2–;	2/4	2–;
U5615-98-136	2-	2-	2	2	;12-	2–;	2	;2-
U5615-98-144 [‡]	2	2/4	2/4	2/4	;1/3+	;2-/4	2/4	2-/4
U5615-98-48	2-/4	2-/4	2	2/4	2-	2-;/4	2/4	;2-
U5616-20-154-7	2-	2-	2	2	2-	2–;	2	2-
U5616-20-47-13	2	2–	2	2	2	;2-	2	2-
U5616-20-47-10	2–	2–	2	2	2	;2-	2	2-
U5616-20-9-13	2-	2	2	2	2	2-;	2	2-
U5616-20-9-15	2–	2	2	2	2	2–;	2	;2-

[†]Race isolates given in parentheses.

both populations markers flanking *Sr22* were identified. In U5615-F_{2:3}, *Sr22* was flanked proximally by *Xcfa2123* at a distance of 0.6 cM and distally at 0.4 cM by cosegregating loci *Xwmc633*, *Xgwm332*, and *Xwmc790*. In the U5616-F_{2:3} population, proximal marker locus *Xcfa2123* was 4.2 cM from *Sr22* and the cosegregating loci *Xwmc633*, *Xgwm332*, and *Xwmc790* were 1.8 cM distal.

Nullisomic and ditelsomic analysis of SSR loci placed Xbarc121, Xcfa2019, Xcfa2123, Xgwm332, Xpsp3094, Xwmc607, Xwmc633, and Xwmc790 on the long arm of chromosome 7A and Xgwm260 on the short arm. Marker Xwmc607 was located on the proximal region between the centromere and the 7AL-1 breakpoint (Fig. 1a). The Xbarc121 locus is located between deletion breakpoints 7AL-10 (fraction length [FL] = 0.49) and 7AL- 21 (FL = 0.74) and the Xcfa2123 locus maps in the submicroscopic region between deletion breakpoints 7AL-8 and 7AL-13, both of which have FL = 0.83. The loci distal to Sr22, Xwmc633, Xgwm332, Xwmc790, Xpsp3094 and Xcfa2019 are located between the breakpoints 7AL-9 (FL = 0.89) and 7AL-15 (FL = 0.99) (Fig. 1a). These data place Sr22 in the deletion interval between the 0.83 and 0.99 breakpoints.

Six SSR loci in the linkage group with *Sr22* including *Xwmc607*, *Xbarc121*, *Xwmc633*, *Xgwm332*, *Xwmc790*, and *Xcfa2019* were evaluated and determined to be polymorphic between Steinwedel and the donor line Sr22Tb (Table 4). These data, combined with our physical mapping of the markers, confirm the result of Paull et al. (1994) that the transfer from *T. boeoticum* in Sr22Tb involves at least 89% of the long arm of chromosome 7A as well as a portion of the short arm.

Although more recombination was observed in the U5616-F_{2:3} population than the U5615-F_{2:3} population, a comparison of the physical location of markers and genetic distances indicate low levels of recombination in

both populations, particularly in the proximal portion of the chromosome. Only 3.5 and 5.6 cM of genetic distance were observed in U5615- $F_{2:3}$ and U5616- $F_{2:3}$ populations, respectively, between markers Xgwm260 and Xcfa2123 that span 83% of the long arm of chromosome 7A plus a portion of the short arm. In the region distal to Sr22, 1.1 and 1.0 cM of genetic distance was observed between SSR markers located in the deletion interval between the breakpoints for 7AL-9 (FL = 0.89) and 7AL-15 (FL = 0.99).

Linkage Analysis of F_{3:4} Recombinant Populations

To develop lines having a smaller amount of T. boeoticum—derived chromatin, additional populations of $F_{3:4}$ families were developed from three recombinant F_2 individuals (U5615-72, U5615-98, and U5616-20; Fig. 2) Linkage

Table 3. Segregation of Sr22 in $F_{2:3}$ and $F_{3:4}$ populations, including number of lines (n), the observed and expected genotypic frequencies, and χ^2 and P values for fit to the 1:2:1 segregation expected for a single dominant gene.

Population	n	Genotype	Observed	χ^2	P
U5615-F _{2:3}	138	Sr22Sr22	35	0.49	0.78
		Sr22sr22	72		
		sr22sr22	31		
U5616- F _{2:3}	139	Sr22Sr22	40	2.61	0.27
		Sr22sr22	72		
		sr22sr22	27		
U5615-72-F _{3:4}	104	Sr22Sr22	21	1.60	0.45
		Sr22sr22	53		
		sr22sr22	30		
U5615-98-F _{3:4}	140	Sr22Sr22	34	0.04	0.98
		Sr22sr22	71		
		sr22sr22	35		
U5616-20-F _{3:4}	152	Sr22Sr22	40	4.63	0.10
		Sr22sr22	64		
		sr22sr22	48		

[‡]Line is heterozygous at marker loci linked to Sr22.

[§]Two or more infection types present on the same plant, with the predominant type given first.

Two scores separated by a slash indicates heterogeneity in infection type, with the predominant phenotype given first.

Table 4. Simple sequence repeat markers used for linkage analysis of *Sr22*, parent allele sizes in base pairs.

Marker	Steinwedel	Sr22Tb	2174	Lakin
Xgwm260	_†	153	163	163
Xwmc607	146	89	143	143
Xbarc121	234	195	217	217
Xcfa2123	_	234	235	241
Xwmc633	260	117	221	229
Xgwm332	184	193	195	275
Xwmc790	152	89	209	100
Xpsp3094	_	228	149	183
Xcfa2019	237	238	217	217

[†]Unavailable data.

analyses of the $F_{3:4}$ populations were performed for loci where the U5615-72, U5615-98, and U5616-20 individuals were heterozygous. In all three $F_{3:4}$ populations, marker orders were consistent with the orders observed in the $F_{2:3}$ populations (Fig. 1). In all populations, *Xwmc633* was closest to *Sr22* with genetic distances ranging from 1.3 to 2.1 cM distal.

The U5615-72- $F_{3:4}$ population consisted of 104 lines that segregated for Sr22, proximal loci Xbarc121, Xcfa2123, Xwmc607, and Xgwm260 as well as distal loci Xgwm332, Xpsp3094, Xwmc633, and Xwmc790. Recombination occurred between the Xcfa2019 locus and Sr22 in the previous generation; thus, the U5615-72- $F_{3:4}$ population was fixed for alleles from parent line 2174 in the region of 7AL distal to the Xcfa2019 locus (Fig. 2). In the related U5615-98 $F_{3:4}$, 140 lines segregated for Sr22 and distal loci Xcfa2019, Xgwm332, Xpsp3094, Xwmc633, and Xwmc790. This population was fixed for alleles from parent line 2174 at the SSR loci proximal to Sr22 (Fig. 2). In both cases, recombination levels were greater (P < 0.01) than observed in the original U5615- $F_{2:3}$ and were similar to the distances observed for the U5616- $F_{2:3}$ and U5616-20- $F_{3:4}$ populations.

Identification of Recombinants

Several recombinant genotypes were identified that carry reduced levels of *T. boeoticum* chromatin associated with *Sr22*. In the U5615-98-F₃₋₄ population that was fixed for

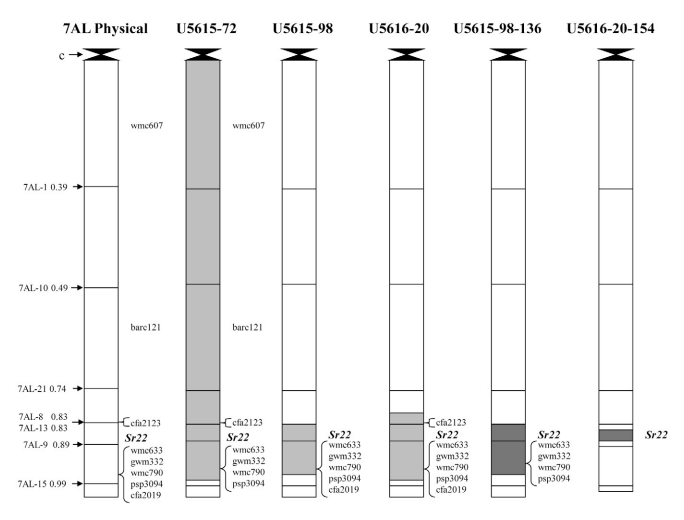


Figure 2. Physical maps of the long arm of chromosome 7A showing *Triticum boeoticum* chromatin in recombinants identified from 2174/Sr22Tb (U5615) and Lakin/Sr22Tb (U5616) F_{2:3} and F_{3:4} populations. Light gray shading represents regions heterozygous for the markers noted to the left of the chromosome. Regions shaded in dark gray represent regions homozygous for *T. boeoticum* chromatin. Deletion breakpoints are indicated as solid lines. Line names and fraction lengths are at left. C indicates the position of the centromere. Marker gwm260 is not shown because it is located on the short arm of chromosome 7A.

alleles from parent line 2174 at marker loci proximal to Sr22, seven genotypes were identified that are resistant to stem rust and in which recombination had occurred between Sr22 and the most distal marker Xcfa2019. Three lines fixed for T. boeoticum alleles at distal loci Xwmc633, Xgwm332, Xwmc790, and Xpsp3094 were homozygous resistant to race RKQQC, of which recombinant U5615-98-136 is shown as example in Fig. 2. Two of these recombinants, U5615-98-136 and U5616-98-48, were evaluated for reaction to nine additional races of stem rust, including TTKSK and TTKST, and exhibited widely effective resistance to all races (Table 2). Four recombinants were heterozygous for Sr22, Xwmc633, Xgwm332, Xwmc790, and *Xpsp3094*, and homozygous for alleles from parent line 2174 at Xcfa2019. Two of these lines (U5615-98-120 and U5615-98-144) were tested and segregated for resistance to diverse races of stem rust (Table 2). These populations can be used to identify new recombinants with further reduced alien chromatin in subsequent generations of inbreeding.

In the U5616-20 population, three genotypes were identified as segregating for resistance to stem rust race RKQQC and homozygous for Lakin alleles at flanking markers *Xcfa2123* and *Xwmc633*, of which recombinant U5616-20-154 is shown as an example in Fig. 2. Plants were self-pollinated and five lines homozygous resistant to all tested races of stem rust were identified (Table 2).

CONCLUSION

Although the stem rust resistance gene Sr22 provides resistance to all known races of Pgt, it has not been widely used in agriculture. This is likely due to detrimental effects associated with introgression from the A^m genome diploid donor species, T. boeoticum. Our data support the hypothesis of Paull et al. (1994) that the proximal regions of both the short and long arm of chromosome 7A are comprised of T. boeoticum chromatin in lines having Sr22 derived from Sr22Tb. Paull et al. (1994) reported that the restriction fragment length polymorphism locus polymorphic between T. boeoticum and hexaploid wheat in the distal region of 7AL, *Xpsr119*, was not transmitted from *T. boeoticum*. They speculated that the terminal region of 7AL in Sr22 hexaploid lines is likely of T. aestivum origin while the proximal region was derived from T. boeoticum. The 13 SSR loci mapped in this study segregated into two unlinked groups, although all markers except Xgwm260 were located on the long arm of 7A. Four of the SSR loci that are located in the most distal region of 7AL were not linked to Sr22 and underwent much greater levels of recombination than observed in the Sr22 region (data not shown). These data agree with the hypothesis that the terminal region of the long arm of 7A in the Sr22Tb donor parent is of *T. aestivum* origin.

Low levels of recombination were observed in the $F_{2:3}$ populations when compared with previous intraspecific SSR maps of chromosome 7A. Somers et al. (2004) reported

the distance between Xgwm260 and Xcfa2019 to be 37 cM. The same interval in our study represented 5.6 cM in the U5615- $F_{2:3}$ population and 12.6 cM in the U5616- $F_{2:3}$ population. Although both populations exhibited suppressed recombination, greater recombination was observed in the $F_{2:3}$ population from the cross Lakin/Sr22Tb than the 2174/Sr22Tb. However, similar levels of recombination in the region were observed in the $F_{3:4}$ populations from both crosses, indicating that the observed reduction in the original 2174/Sr22Tb $F_{2:3}$ may have been specific to that population.

The and McIntosh (1975) reported reduced transmission of *Sr22* in pooled populations of lines with *Sr22* from both *T. monococcum* and *T. boeoticum*. In this study, segregation for 7AL alleles including *Sr22* did not deviate significantly from the expected 1:2:1 ratio in all populations.

Previously, the SSR loci *Xcfa2123* and *Xcfa2019* were identified as flanking markers for *Sr22* (Khan et al., 2005). These same markers were polymorphic in both of our populations and were found to flank *Sr22*. Reduced levels of recombination associated with *Sr22* were identified by Khan et al. (2005) in a single F₄–derived, single seed descent population, on which mapping was done only on homozygous resistant and susceptible lines. In the present study, all individuals in multiple populations were genotyped with codominant markers allowing for more accurate estimates of genetic distances. Mapping in the F_{2:3} allowed for the identification of critical recombinants with reduced *T. boeoticum* chromatin. From these recombinants, additional mapping populations were made and recombinants with even further reduced alien chromatin were identified.

The recombinant lines carrying reduced T. boeoticum chromatin identified in this study will be useful in breeding programs. Using these lines as donor parents should help to meliorate the negative effects associated with Sr22. These recombinant lines are being backcrossed to hard and soft winter wheat cultivars to produce populations and NILs for which the effects of the introgression on agronomic performance can be evaluated. The marker loci documented in this study will prove useful in the marker-assisted selection of Sr22. The most closely linked marker locus to Sr22 across all populations was the distal locus Xwmc633, while the proximal flanking marker Xcfa2123 could also be used but greater recombination between this locus and Sr22 was observed. Markers identified in this study can be used for selection among the progeny of the recombinant lines from the U5615-98 population to further reduce the size of the T. boeoticum chromosome segment. Homozygous resistant F_{4.5} progeny have been identified from each of the U5616-20-derived recombinant plants. The use of marker-assisted selection in progeny from crosses with the U5616-20 recombinants will require the development of markers more closely linked to Sr22. Recombinants with reduced T. boeoticum chromatin are available on request for development of wheat cultivars having resistance to stem rust.

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References

- Allard, R. 1956. Formulas and tables to facilitate the calculation of recombination values in heredity. Hilgardia 24:235–278.
- Dubcovsky, J., M.C. Luo, and J. Dvorak. 1995. Differentiation between homoeologous chromosomes 1A of wheat and 1A(M) of *Triticum monococcum* and its recognition by the wheat *Ph1* locus. Proc. Natl. Acad. Sci. USA 92:6645–6649.
- Endo, T.R., and B.S. Gill. 1996. The deletion stocks of wheat. J. Hered. 87:295–307.
- Gerechter-Amati, Z.K., I. Wahl, A. Vardi, and D. Zohary. 1971. Transfer of stem rust seedling resistance from wild diploid einkorn to tetraploid durum wheat by means of a triploid hybrid bridge. Euphytica 20:281–285.
- Jin, Y., R.P. Singh, R.W. Ward, R. Wanyera, M. Kinyua, P. Njau, and Z.A. Pretorius. 2007. Characterization of seedling infection types and adult plant infection responses of monogenic Sr gene lines to race TTKS of *Puccinia graminis* f. sp. tritici. Plant Dis. 91:1096–1099.
- Jin, Y., L.J. Szabo, Z.A. Pretorius, R.P. Singh, R. Ward, and T. Fetch, Jr. 2008. Detection of virulence to resistance gene Sr24 within race TTKS of Puccinia graminis f. sp. tritici. Plant Dis. 92:923–926.
- Kerber, E.R., and P.L. Dyck. 1973. Inheritance of stem rust resistance transferred from diploid wheat (*Triticum monococcum*) to tetraploid and hexaploid wheat and chromosome location of gene involved. Can. J. Genet. Cytol. 15:397–409.
- Khan, R.R., H.S. Bariana, B.B. Dholakia, S.V. Naik, M.D. Lagu, A.J. Rathjen, and V.S. Gupta. 2005. Molecular mapping of stem and leaf rust resistance in wheat. Theor. Appl. Genet. 111:846–850.
- Lander, E.S., P. Green, J. Abrahamson, A. Barlow, M.J. Daly, S.E. Lincoln, and L.A. Newberg. 1987. MAPMAKER: An interactive computer package for constructing primary genetic linkage maps of experimental and natural populations. Genomics 2:174–181.
- Luo, M.C., Z.L. Yang, R.S. Kota, and J. Dvorak. 2000. Recombination of chromosomes 3A(m) and 5A(m) of *Triticum monococcum* with homeologous chromosomes 3A and 5A of wheat: The distribution of recombination across chromosomes. Genetics 154:1301–1308.
- McFadden, E.S. 1930. A successful transfer of emmer characteristics to vulgare wheat. J. Am. Soc. Agron. 22:1020–1034.
- Miranda, L.M., L. Perugini, G. Srnic, G. Brown-Guedira, D. Marshall, S. Leath, and J.P. Murphy. 2007. Genetic mapping of a *Triticum monococcum* derived powdery mildew derived resistance gene in common wheat. Crop Sci. 47:2323–2329.
- Nyquist, W.E. 1962. Differential fertilization in the inheritance of stem rust resistance in hybrids involving a common wheat strain derived from *Triticum timopheevi*. Genetics 47:1109–1124.
- Pallotta, M.A., P. Warner, R.L. Fox, H. Kuchel, S.J. Jefferies, and P. Langridge. 2003. Marker assisted wheat breeding in the southern region of Australia. p. 789–791. *In* N.E. Pogna et al. (ed.)
 Proc. 10th Int. Wheat Genet. Symp., Paestum, Italy. 1–6 Sept.

- 2003. Instituto Sperimentale per la Cearealicoltura, Rome.
- Paull, J.G., M.A. Pallotta, P. Langridge, and T.T. The. 1994. RFLP markers associated with *Sr22* and recombination between chromosome 7A of bread wheat and the diploid species *Triticum boeoticum*. Theor. Appl. Genet. 89:1039–1045.
- Perugini, L.D., J.P. Murphy, D. Marshall, and G. Brown-Guedira. 2007. *Pm37*, a new broadly effective powdery mildew resistance gene from *Triticum timopheevii*. Theor. Appl. Genet. 11:417–425.
- Riley, R., and V. Chapman. 1958. Genetic control of the cytologically diploid behavior of hexaploid wheat. Nature 182:713–715.
- Roelfs, A.J., and J.W. Martens. 1988. An international system of nomenclature for *Puccinia graminis* f. sp. tritici. Phytopathology 78:526–533.
- Sears, E.R. 1954. The aneuploids of common wheat. Univ. Mo. Agric. Exp. Stn. Bull. 572:1Y58.
- Sears, E.R. 1973. *Agropyron-wheat* transfers induced by homoeologous pairing. p. 191–199. *In* E.R. Sears and L.M.S. Sears (ed.) Proc. 4th Int. Wheat Genet. Symp., Columbia, MO. 6–11 Aug. 1973. Univ. of Missouri, Columbia.
- Sears, E.R., and L.M.S. Sears. 1978. The telocentric chromosomes of common wheat. p. 389–407. *In* S. Ramanujam (ed.) Proc. 5th Int. Wheat Genet. Symp., New Delhi, India. 23–28 Feb. 1978. Indian Soc. Genetics and Plant Breeding, New Delhi.
- Singh, R.P., D.P. Hodson, J. Huerta-Espino, Y. Jin, P. Njau, R. Wanyera, S.A. Herrera Foessel, and R.W. Ward. 2008. Will stem rust destroy the world's wheat crop? p. 271–309 In D.L. Sparks (ed.) Advances in agronomy, Vol. 98. Elsevier Academic Press Inc., San Diego.
- Somers, D.J., P. Issac, and K. Edwards. 2004. A high density microsatellite map for bread wheat (*Triticum aestivum* L.). Theor. Appl. Genet. 109:1105–1114.
- Sourdille, P., B. Grandon, V. Chiquet, N. Nicot, D. Somers, A. Murigneux, and M. Bernard. 2005. Wheat Genoplante SSR mapping data release: A new set of markers and comprehensive genetic and physical mapping data. Available at http://wheat.pw.usda.gov/ggpages/SSR.club/GeneticPhysical/ (verified 24 June 2010).
- Stakman, E.C., D.M. Steward, and W.Q. Loegering. 1962. Identification of physiologic races of *Puccinia graminis* var. tritici. U.S. Dep. Agric. Res. Serv. E-617.
- The, T.T. 1973. Chromosome location of genes conditioning stem rust resistance transferred from diploid to hexaploid wheat. Nat. New Biol. 241:256.
- The, T.T., B.D.H. Latter, R.A. McIntosh, F.W. Ellison, P.S. Brennan, J. Fisher, G.J. Hollamby, A.J. Rathjen, and R.E. Wilson. 1988. Grain yields of near-isogenic lines with added genes for stem rust resistance. p. 901–906. *In* T.E. Miller and R.M.D. Koebner (ed.) Proc. 7th Int. Wheat Genet. Symp., Cambridge, UK. 13–19 July 1988. Inst. of Plant Sci. Res., Cambridge.
- The, T.T., and R.A. McIntosh. 1975. Cytogenetical studies in wheat 8. Telocentric mapping and linkage studies involving *Sr22* and other genes on chromosome 7AL. Aust. J. Biol. Sci. 28:531–538.
- Zeller, F.J. 1973. 1B/1R wheat-rye chromosome substitutions and translocations. p. 209–222. *In* E.R. Sears and L.M.S. Sears (ed.) Proc. 4th Int. Wheat Genet. Symp., Columbia, MO. 6–11 Aug. 1973. Univ. of Missouri, Columbia.